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FILE 'MEDLINE, EMBASE, BIOSIS, BIOTECHDS, SCISEARCH, HCAPLUS, NTIS,
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L1
      10975 S ADENYLATE KINASE?
           2615 S HUMAN AND L1
L2
         613747 S MITOCHONDR?
L3
            255 S L2 AND L3
L4
        5929362 S CLON? OR EXPRESS? OR RECOMBINANT
L5
             59 S L4 AND L5
L6
             32 DUP REM L6 (27 DUPLICATES REMOVED)
ь7
Г8
              8 S "HMAK"
L9
              3 DUP REM L8 (5 DUPLICATES REMOVED)
                E HILLMAN J L/AU
L10
            454 S E3
              1 S L4 AND L10
L11
                E SHAH P/AU
           1409 S E3
L12
              1 S L4 AND L12
L13
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                PHARMAMarketLetter(PHARMAML) - new on STN
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        Aug 08
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                now available on STN
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NEWS 18 Dec 17 Adis Clinical Trials Insight now available on STN
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                CANCERLIT is no longer being updated
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                structures available in REGISTRY
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        Apr 11 Display formats in DGENE enhanced
NEWS 32
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                MEDLINE Reload
NEWS 33
        Apr 17
                Polymer searching in REGISTRY enhanced
                Indexing from 1947 to 1956 being added to records in CA/CAPLUS
NEWS 34
        Apr 21
NEWS 35
                New current-awareness alert (SDI) frequency in
        Apr 21
                WPIDS/WPINDEX/WPIX
NEWS 36
        Apr 28
                RDISCLOSURE now available on STN
                Pharmacokinetic information and systematic chemical names
NEWS 37
        May 05
                 added to PHAR
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NEWS EXPRESS April 4 CURRENT WINDOWS VERSION IS V6.01a, CURRENT MACINTOSH VERSION IS V6.0b(ENG) AND V6.0Jb(JP), AND CURRENT DISCOVER FILE IS DATED 01 APRIL 2003

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0.21

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=> s human and l1 L2 2615 HUMAN AND L1

=> s mitochondr?

=> s 12 and 13

255 L2 AND L3

=> s clon? or express? or recombinant

3 FILES SEARCHED...

5929362 CLON? OR EXPRESS? OR RECOMBINANT

=> s 14 and 15

59 L4 AND L5

=> dup rem 16

PROCESSING COMPLETED FOR L6

32 DUP REM L6 (27 DUPLICATES REMOVED)

=> d 1-32 ibib ab

ANSWER 1 OF 32 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

2002:637880 HCAPLUS

DOCUMENT NUMBER:

137:179893

TITLE:

Methods for identifying compounds that inhibit or reduce PTP1B (protein tyrosine phosphatase 1B)

INVENTOR(S):

Zinker, Bradley A.; Trevillyan, James M.; Jirousek, Michael R.; Rondinone, Christina M.; Cowsert, Lex M.; Wyatt, Jacqueline; Monia, Brett P.; Butler, Madeline

M.; Waring, Jeffrey French

PATENT ASSIGNEE(S):

Abbott Laboratories, USA; Isis Pharmaceuticals, Inc.

SOURCE:

PCT Int. Appl., 72 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

APPLICATION NO. DATE PATENT NO. KIND DATE \_\_\_\_\_ WO 2002-US4194 WO 2002064840 **A**2 20020822 20020213

W: CA, JP, MX

RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR

PRIORITY APPLN. INFO.:

US 2001-268399P P 20010213 US 2002-74194 A 20020212

The present invention relates to methods for identifying compds. that AB inhibit PTP1B (protein tyrosine phosphatase 1B) mRNA and protein expression in insulin resistant obese non-human mammals. The present invention relates to biol. markers for PTP1B inhibition or redn. Specifically, the present invention relates to methods for measuring the downregulation of the p85.alpha. regulatory subunit of phosphatidylinositol-3-kinase and the upregulation of p55.alpha. and/or p50.alpha. isoforms in response to in vivo inhibition or redn. of PTP1B in insulin resistant mammals. Moreover, the present invention relates to an in vivo marker for pharmacodynamic measurements and mechanism of action detns. of small mol. drugs which inhibit or reduce PTP1B activity. Finally, the present invention also provides a method to screen agents for activity that down modulates p85.alpha. and upregulates phosphatidylinositol-3-kinase p85.alpha. isoforms as drugs for the treatment of type 2 diabetes.

ANSWER 2 OF 32 HCAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER: 2002:937303 HCAPLUS

DOCUMENT NUMBER: 138:20443

TITLE: Endocrine disruptor screening using DNA chips of

endocrine disruptor-responsive genes

INVENTOR(S): Kondo, Akihiro; Takeda, Takeshi; Mizutani, Shigetoshi;

Tsujimoto, Yoshimasa; Takashima, Ryokichi; Enoki,

Yuki; Kato, Ikunoshin

PATENT ASSIGNEE(S): Takara Bio Inc., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 386 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE		APPLICATION N	0.	DATE
JP 2002355079	A2	20021210		JP 2002-69354		20020313
PRIORITY APPLN. INFO.		JP	2001-73183	Α	20010314	
			JP	2001-74993	Α	20010315
			JΡ	2001-102519	A	20010330

AB A method and kit for detecting endocrine-disrupting chems. using DNA microarrays are claimed. The method comprises prepg. a nucleic acid sample contg. mRNAs or cDNAs originating in cells, tissues, or organisms which have been brought into contact with a sample contg. the endocrine disruptor. The nucleic acid sample is hybridized with DNA microarrays having genes affected by the endocrine disruptor or DNA fragments originating in these genes have been fixed. The results obtained are then compared with the results obtained with the control sample to select the gene affected by the endocrine disruptor. Genes whose expression is altered by tri-Bu tin, 4-octaphenol, 4-nonylphenol, di-N-Bu phthalate, dichlorohexyl phthalate, octachlorostyrene, benzophenone, diethylhexyl phthalate, diethylstilbestrol (DES), and 17-.beta. estradiol (E2), were found in mice by DNA chip anal.

L7 ANSWER 3 OF 32 EMBASE COPYRIGHT 2003 ELSEVIER SCI. B.V.

ACCESSION NUMBER: 2002449003 EMBASE

TITLE: Old and new determinants in the regulation of energy

expenditure.

AUTHOR: Russell A.P.; Giacobino J.P.

CORPORATE SOURCE: Prof. J.P. Giacobino, Departement de Biochimie Medicale,

Centre Medical Universitaire, 1 rue Michel-Servet, 1211 Geneve 4, Switzerland. Jean-Paul.Giacobino@medecine.unige.c

h

SOURCE: Journal of Endocrinological Investigation, (2002) 25/10

(862-866). Refs: 55

ISSN: 0391-4097 CODEN: JEIND7

COUNTRY:

Italy

DOCUMENT TYPE: Journal; General Review FILE SEGMENT: 003 Endocrinology

029 Clinical Biochemistry

005 General Pathology and Pathological Anatomy

030 Pharmacology

037 Drug Literature Index

LANGUAGE: English SUMMARY LANGUAGE: English

AB Bw gain is controlled by energy intake on one hand and expenditure on the other. The components of energy expenditure are basal metabolism, exercise induced thermogenesis and adaptive thermogenesis. In this short review we shall discuss the main determinants of adaptive thermogenesis.

.COPYRGT.2002, Editrice Kurtis.

L7 ANSWER 4 OF 32 EMBASE COPYRIGHT 2003 ELSEVIER SCI. B.V.

ACCESSION NUMBER: 2002446521 EMBASE

TITLE: Frontiers in research on parasitic protozoa.

AUTHOR: Gibson W.; Miles M.

CORPORATE SOURCE: W. Gibson, School of Biological Sciences, University of

Bristol, Woodland Road, Bristol BS8 1UG, United Kingdom.

w.gibson@bristol.ac.uk

SOURCE: Trends in Parasitology, (1 Dec 2002) 18/12 (521-522).

Refs: 2

ISSN: 1471-4922 CODEN: TPRACT

PUBLISHER IDENT.: S 1471-4922(02)02416-9

COUNTRY: United Kingdom

DOCUMENT TYPE: Journal; Conference Article

FILE SEGMENT: 004 Microbiology

037 Drug Literature Index

LANGUAGE: English

L7 ANSWER 5 OF 32 BIOTECHDS COPYRIGHT 2003 THOMSON DERWENT AND ISI

ACCESSION NUMBER: 2002-00258 BIOTECHDS

TITLE: New antibody against human mitochondria

adenylate-kinase isozyme 2 or isozyme 3,

for detecting the isozymes in a detection sample to diagnose cardiac diseases such as myocardial infarction and angina

pectoris;

monoclonal antibody, hybridoma cell culture and detection

marker useful in disease diagnosis

AUTHOR: Cho K S; Lee S M

PATENT ASSIGNEE: Kim H J

LOCATION: Ansan, Korea.

PATENT INFO: WO 2001058482 16 Aug 2001
APPLICATION INFO: WO 2000-KR882 10 Aug 2000
PRIORITY INFO: KR 2000-5808 8 Feb 2000

DOCUMENT TYPE: Patent LANGUAGE: English

OTHER SOURCE: WPI: 2001-522438 [57]

AB An antibody (I) specific to human mitochondria

adenylate-kinase (AK) isozymes AK2 or AK3 or their

portion, is claimed. (I) is produced in an animal species and has a

reactivity with the immunogen which includes a human

mitochondria adenylate-kinase isozyme or its

portion. Also claimed are: an immunological formulation (II) for diagnosing cardiac disease containing (I) and a detection marker; and a diagnostic kit (III) for cardiac disease containing a carrier and (I) which is coupled with a detection marker. (I) is useful for detecting a

human mitochondrial adenylate-kinase

isozyme (AK2) or(AK3) in a detection sample. An immunological formulation (II) for diagnosing cardiac disease containing (I) and a detection marker is useful for detecting adenylate-

kinase isozyme in a biological sample. (I) is useful for

diagnosing cardiac disease such as myocardial infarction, angina pectoris. (II) and a diagnostic kit (III) are also useful for

diagnosing cardiac disease. (56pp)

L7 ANSWER 6 OF 32 HCAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER: 2001:597830 HCAPLUS

DOCUMENT NUMBER: 135

135:194482

TITLE:

Anti-human mitochondrial

adenylate kinase isozyme antibody,

diagnostic formulation and diagnostic kit for cardiac

disease

INVENTOR(S):

Kim, Hyo Joon; Cho, Key Seung; Lee, Sang Min

PATENT ASSIGNEE(S):

S. Korea

SOURCE:

PCT Int. Appl., 56 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

DATE PATENT NO. KIND DATE APPLICATION NO. A1 20010816 WO 2000-KR882 WO 2001058482 20000810

W: AU, BR, CA, CN, DE, JP, RU, US

RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,

PT, SE

20020102 EP 2000-955110 20000810 EP 1165133 A1

AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,

IE, FI

BR 2000010712 Α 20020205 BR 2000-10712 20000810 A 20000208 PRIORITY APPLN. INFO.: KR 2000-5808

WO 2000-KR882 W 20000810

The present invention relates to an immunol. formulation and a diagnostic AB kit for cardiac disease using human mitochondrial adenylate kinase isoenzymes. The present invention provides an immunol. formulation and a diagnostic kit for cardiac disease, which are featured by using mitochondrial adenylate kinase isoenzymes which exist in a myocardiac cell among muscle cells, but not in a skeletal muscular cell, as a diagnostic marker for cardiac disease and which enable more correct and easy diagnosis of

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 7 OF 32 MEDLINE

cardiac disease.

DUPLICATE 1

ACCESSION NUMBER:

2001439872

4

MEDLINE

DOCUMENT NUMBER:

TITLE:

21378190 PubMed ID: 11485571

Structure and expression of human mitochondrial adenylate kinase

targeted to the mitochondrial matrix.

Noma T; Fujisawa K; Yamashiro Y; Shinohara M; Nakazawa A; AUTHOR:

Gondo T; Ishihara T; Yoshinobu K

Department of Biochemistry, Yamaguchi University School of CORPORATE SOURCE:

Medicine, 1-1-1 Minami-Kogushi, Ube, Yamaguchi 755-8505,

Japan.. tnoma@po.cc.yamaguchi-u.ac.jp

BIOCHEMICAL JOURNAL, (2001 Aug 15) 358 (Pt 1) 225-32. SOURCE:

Journal code: 2984726R. ISSN: 0264-6021.

PUB. COUNTRY: DOCUMENT TYPE:

England: United Kingdom Journal; Article; (JOURNAL ARTICLE)

English LANGUAGE:

FILE SEGMENT: Priority Journals GENBANK-AB021870 OTHER SOURCE:

ENTRY MONTH:

200109

ENTRY DATE: Entered STN: 20010924

Last Updated on STN: 20010924 Entered Medline: 20010920

AB The previously isolated cDNA encoding human adenylate kinase (AK) isozyme 3 was recently renamed AK4. Consequently, human AK3 cDNA remains to be identified and we have little information about the functional relationship between human AK3 and AK4. In pursuit of the physiological roles of both the AK3 and AK4 proteins, we first isolated an authentic human AK3 cDNA and compared their expression. Nucleotide sequencing revealed that the cDNA encoded a 227-amino-acid protein, with a deduced molecular mass of 25.6 kDa, that shares greater homology with the AK3 cDNAs isolated from bovine and rat than that from human. We named the isolated cDNA AK3. Northern-blot analysis revealed that AK3 mRNA was present in all tissues examined, and was highly expressed in heart, skeletal muscle and liver, moderately expressed in pancreas and kidney, and weakly expressed in placenta, brain and lung. On the other hand, we found that human AK4 mRNA was highly expressed in kidney, moderately expressed in heart and liver and weakly expressed in brain. Western-blot analysis demonstrated expression profiles of AK3 and AK4 that were similar to their mRNA expression patterns in each tissue. Over expression of AK3, but not AK4, in both Escherichia coli CV2, a temperature-sensitive AK mutant, and a human embryonic kidney-derived cell line, HEK-293, not only produced significant GTP:AMP phosphotransferase (AK3) activity, but also complemented the CV2 cells at 42 degrees C. Subcellular and submitochondrial fractionation analysis demonstrated that both AK3 and AK4 are localized in the mitochondrial matrix.

ANSWER 8 OF 32 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC.

ACCESSION NUMBER: DOCUMENT NUMBER:

2002:288316 BIOSIS PREV200200288316

TITLE:

Hemodynamic unloading by ventricular assist devices has no beneficial effect for the inflammation-associated apoptotic

pathway in human terminally failing myocardium.

AUTHOR(S):

Scheubel, Robert Johannes (1); Bartling, Babett (1); Stein,

Susanne; Darmer, Dorothea; Holtz, Juergen; Silber,

Rolf-Edgar

CORPORATE SOURCE:

(1) Clin fuer Herz- und Thoraxchirurgie, Halle/Saale

Germany

SOURCE:

Circulation, (October 23, 2001) Vol. 104, No. 17 Supplement, pp. II.713. http://circ.ahajournals.org/.

print.

Meeting Info.: Scientific Sessions 2001 of the American Heart Association Anaheim, California, USA November 11-14,

2001

ISSN: 0009-7322.

DOCUMENT TYPE: LANGUAGE:

Conference English

ANSWER 9 OF 32 HCAPLUS COPYRIGHT 2003 ACS · 2000:810680 HCAPLUS

ACCESSION NUMBER: DOCUMENT NUMBER:

133:345587

TITLE:

Protein and cDNA sequences of a novel human

Mitochondria adenylate

kinase GTP3P and uses thereof

INVENTOR(S):

Yu, Long; Zhao, Yong; Bi, Anding; Gao, Jie; Zhao,

Shouyuan

PATENT ASSIGNEE(S):

Fudan Gene Engineering Co., Ltd., Xinhuangpu,

Shanghai, Peop. Rep. China

SOURCE:

Faming Zhuanli Shenqing Gongkai Shuomingshu, 20 pp.

CODEN: CNXXEV

DOCUMENT TYPE:

Patent

LANGUAGE:

Chinese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

DATE APPLICATION NO. PATENT NO. KIND CN 1249340 Α 20000405 CN 1998-119439 19980928 PRIORITY APPLN. INFO.: CN 1998-119439

The invention provides protein and cDNA sequences of a novel human Mitochondria adenylate kinase GTP3P which is

belived to be a GTP-AMP transphosphorylase. The invention also relates to

constructing adenylate kinase GTP3P expression cassette to producing recombinant adenylate kinase GTP3P using E.coli cells or eukaryotic cells. invention further relates to the uses of adenylate kinase GTP3P.

ANSWER 10 OF 32 SCISEARCH COPYRIGHT 2003 THOMSON ISI

2000:361124 SCISEARCH ACCESSION NUMBER:

THE GENUINE ARTICLE: 311BC

Cellular phosphorylation of 2',3'-dideoxyadenosine-5'-TITLE:

monophosphate, a key intermediate in the activation of the

antiviral agent DDI, in huhlan peripheral blood

mononuclear cells

Robbins B L (Reprint); Greenshaw J; Fridland A AUTHOR:

CORPORATE SOURCE: ST JUDE CHILDRENS HOSP, DEPT INFECT DIS, 332 N LAUDERDALE

ST, MEMPHIS, TN 38105 (Reprint)

COUNTRY OF AUTHOR: USA

SOURCE: NUCLEOSIDES NUCLEOTIDES & NUCLEIC ACIDS, (MAY 2000) Vol.

19, No. 1-2, pp. 405-413.

Publisher: MARCEL DEKKER INC, 270 MADISON AVE, NEW YORK,

NY 10016.

ISSN: 1525-7770.

DOCUMENT TYPE:

Article; Journal

FILE SEGMENT:

LIFE

LANGUAGE:

English

REFERENCE COUNT:

20

\*ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS\*

2',3'-dideoxyadenosine 5-monophosphate (ddAMP), is a key intermediate AB in the metabolism of the antiviral agent 2',3'-dideoxyinosine (ddI) to its active triphosphate derivative, 2',3'-dideoxyadenosine-5'-triphosphate (ddATP). The potential role of adenylate kinase in the phosphorylation of ddAMP was studied in human peripheral blood mononuclear cells (PBMC) and a human T cell line, CEMss. Subcellular distribution, sulfhydryl inhibitor, and substrate specificity studies support the hypothesis that the mitochondrial adenylate kinase (AK2) is a major route of cellular

activation of these compounds in human lymphocytes. ANSWER 11 OF 32 MEDLINE DUPLICATE 2

ACCESSION NUMBER: 2000246295 MEDLINE

20246295 PubMed ID: 10786623

DOCUMENT NUMBER:

TITLE: cDNA cloning and chromosomal mapping of the gene

encoding adenylate kinase 2 from

Drosophila melanogaster.

Noma T; Murakami R; Yamashiro Y; Fujisawa K; Inouye S; AUTHOR:

Nakazawa A

CORPORATE SOURCE: Department of Biochemistry, Yamaguchi University School of

Medicine, Ube, Japan.. tnoma@po.cc.yamaguchi-u.ac.jp

BIOCHIMICA ET BIOPHYSICA ACTA, (2000 Jan 31) 1490 (1-2) SOURCE:

109-14.

Journal code: 0217513. ISSN: 0006-3002.

PUB. COUNTRY: Netherlands

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

OTHER SOURCE: GENBANK-AB009996; GENBANK-AC004642

ENTRY MONTH: 200005

ENTRY DATE: Entered STN: 20000606

> Last Updated on STN: 20000606 Entered Medline: 20000524

AB As a step toward understanding of the role of adenylate kinase (AK) in energy metabolism, we analyzed this enzyme in

Drosophila melanogaster. The enzyme activities of all three AK isozymes were determined in cell-free extracts of flies, and their proteins were detected by Western blot analysis using polyclonal antibodies against the mammalian isozymes. A cDNA encoding adenylate kinase was isolated from D. melanogaster cDNA library. The cDNA encodes a 240-amino acid protein, which shows high similarity to bovine, human and rat AK2, and hence was named DAK2. Preliminary subcellular fractionation analysis indicated that DAK2 is localized in both cytoplasm and mitochondria. In situ hybridization to salivary gland polytene chromosomes revealed that the Dak2 gene is located at 60B on the right arm of the second chromosome.

ANSWER 12 OF 32 SCISEARCH COPYRIGHT 2003 THOMSON ISI

ACCESSION NUMBER:

2000:126507 SCISEARCH

THE GENUINE ARTICLE: 282KY

TITLE:

cDNA cloning and chromosomal mapping of the gene

encoding adenylate kinase 2 from

Drosophila melanogaster

AUTHOR:

Noma T (Reprint); Murakami R; Yamashiro Y; Fujisawa K;

Inouye S; Nakazawa A

CORPORATE SOURCE:

YAMAGUCHI UNIV, SCH MED, DEPT BIOCHEM, YAMAGUCHI 7558505, JAPAN (Reprint); YAMAGUCHI UNIV, FAC SCI, DEPT PHYS BIOL &

INFORMAT, YAMAGUCHI 7538512, JAPAN

COUNTRY OF AUTHOR:

SOURCE:

BIOCHIMICA ET BIOPHYSICA ACTA-GENE STRUCTURE AND

EXPRESSION, (31 JAN 2000) Vol. 1490, No. 1-2, pp. 109-114.

Publisher: ELSEVIER SCIENCE BV, PO BOX 211, 1000 AE

AMSTERDAM, NETHERLANDS.

ISSN: 0167-4781.

DOCUMENT TYPE:

Article; Journal

FILE SEGMENT: LANGUAGE:

LIFE English

REFERENCE COUNT: 32

\*ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS\*

AB As a step toward understanding of the role of adenylate kinase (AK) in energy metabolism, we analyzed this enzyme in Drosophila melanogaster. The enzyme activities of all three AK isozymes were determined in cell-free extracts of flies, and their proteins were detected by Western blot analysis using polyclonal antibodies against the mammalian isozymes. A cDNA encoding adenylate kinase was isolated from D, melanogaster cDNA library. The cDNA encodes a 240-amino acid protein, which shows high similarity to bovine, human and rat AK2, and hence was named DAK2. Preliminary subcellular fractionation analysis indicated that DAK2 is localized in both cytoplasm and mitochondria. In situ hybridization to salivary gland polytene chromosomes revealed that the Dak2 gene is located at 60B on the right arm of the second chromosome. (C) 2000 Elsevier Science B.V. All rights reserved.

ANSWER 13 OF 32 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

2001:782966 HCAPLUS

DOCUMENT NUMBER:

136:322434

TITLE:

Expression of mRNAs encoding

adenylate kinase isozymes 1, 2, 3,

and 4 in mouse tissues and during neuronal

differentiation of P19 embryonal carcinoma cells

Yamashiro, Yasuhiro

CORPORATE SOURCE:

Department of Biochemistry, Yamaguchi University

School of Medicine, Yamaguchi, 755-8505, Japan Bulletin of the Yamaguchi Medical School (2000),

SOURCE:

AUTHOR(S):

47(3-4), 55-68

CODEN: BYMSAN; ISSN: 0513-1812

PUBLISHER: Yamaguchi University, School of Medicine

DOCUMENT TYPE: Journal LANGUAGE: English

AB The authors cloned cDNAs encoding four adenylate

kinase (AK) isoenzymes from a mouse kidney cDNA library. The AK1, AK2, AK3, and AK4 cDNAs encode the 194-, 232-, 227-, and 223-amino acid proteins, resp. AK4 is a recently isolated gene that is highly homologous to the reported human AK3. Northern blot anal. and reverse transcription-polymerase chain reaction anal. revealed that AK1 mRNA was

predominantly expressed in skeletal muscle, heart, and testis;

AK2 mRNA in liver, heart, kidney, and testis; AK3 mRNA almost uniformly in all tissues examd.; and AK4 mRNA prominently in kidney. Subcellular and submitochondrial fractionation anal. suggested that AK4 was localized in

the mitochondrial matrix. Further, the authors found a 76-fold induction of AK1 mRNA expression concomitant with a 53-fold induction of NeuroD expression during retinoic acid-induced

neuronal differentiation of P19 embryonic carcinoma cell. AK2 and AK3

mRNA expression was increased by 4- to 6-fold during

differentiation, whereas AK4 transcription was first down-regulated and subsequently returned to the original level. These data on AK isoenzyme gene **expression** may provide basic information for prodn. and

evaluation of transgenic mice as well as knockout mice to further understand the physiol. role of AK isoenzymes.

REFERENCE COUNT: 43 THERE ARE 43 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 14 OF 32 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC.

ACCESSION NUMBER: 2000:290852 BIOSIS DOCUMENT NUMBER: PREV200000290852

TITLE: Mitochondrial adenylate kinase

AUTHOR(S): Hillman, Jennifer L.; Shah, Purvi

ASSIGNEE: Incyte Pharmaceuticals, Inc.

PATENT INFORMATION: US 6001624 December 14, 1999

SOURCE: Official Gazette of the United States Patent and Trademark

Office Patents, (Dec. 14, 1999) Vol. 1229, No. 2, pp. No.

pagination. e-file. ISSN: 0098-1133.

DOCUMENT TYPE: Patent LANGUAGE: English

AB The present invention provides a human mitochondrial adenylate kinase (HMAK) and polynucleotides which encode HMAK. The invention also provides expression vectors, host

cells, agonists, antisense molecules, antibodies, or antagonists. The invention also provides methods for treating disorders associated with expression of HMAK.

L7 ANSWER 15 OF 32 MEDLINE DUPLICATE 3

ACCESSION NUMBER: 1999221639 MEDLINE

DOCUMENT NUMBER: 99221639 PubMed ID: 10205158

TITLE: Presence of a pre-apoptotic complex of pro-caspase-3, Hsp60

and Hsp10 in the mitochondrial fraction of jurkat

cells.

AUTHOR: Samali A; Cai J; Zhivotovsky B; Jones D P; Orrenius S CORPORATE SOURCE: Institute of Environmental Medicine, Division of

Toxicology, Karolinska Institutet, Box 210, S-171 77,

Stockholm, Sweden.. afshin.samali@imm.ki.se

CONTRACT NUMBER: ES09047 (NIEHS)

SOURCE: EMBO JOURNAL, (1999 Apr 15) 18 (8) 2040-8. Journal code: 8208664. ISSN: 0261-4189.

PUB. COUNTRY: ENGLAND: United Kingdom

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199906

ENTRY DATE: Entered STN: 19990628

Last Updated on STN: 19990628 Entered Medline: 19990611

Activation of pro-caspase-3 is a central event in the execution phase of AB apoptosis and appears to serve as the convergence point of different apoptotic signaling pathways. Recently, mitochondria were found to play a central role in apoptosis through release of cytochrome c and activation of caspases. Moreover, a sub-population of pro-caspase-3 has been found to be localized to this organelle. In the present study, we demonstrate that pro-caspase-3 is present in the mitochondrial fraction of Jurkat T cells in a complex with the chaperone proteins Hsp60 and Hsp10. Induction of apoptosis with staurosporine led to the activation of mitochondrial pro-caspase-3 and its dissociation from the Hsps which were released from mitochondria. The release of Hsps occurred simultaneously with the release of other mitochondrial intermembrane space proteins including cytochrome c and adenylate kinase, prior to a loss of mitochondrial transmembrane potential. In in vitro systems, recombinant Hsp60 and Hsp10 accelerated the activation of pro-caspase-3 by cytochrome c and dATP in an ATP-dependent manner, consistent with their function as chaperones. This finding suggests that the release of mitochondrial Hsps may also accelerate caspase activation in the cytoplasm of intact cells.

L7 ANSWER 16 OF 32 BIOTECHDS COPYRIGHT 2003 THOMSON DERWENT AND ISI

ACCESSION NUMBER: 1999-00127 BIOTECHDS

TITLE: Human mitochondrial adenylate-

kinase, HMAK;

sense, antisense sequence, antibody, agonist and

antagonist used for cancer, neurological and immunological

disorder diagnosis and therapy

AUTHOR: Hillman J L; Shah P

PATENT ASSIGNEE: Incyte-Pharm.

LOCATION: Palo Alto, CA, USA. PATENT INFO: WO 9844124 8 Oct 1998

APPLICATION INFO: WO 1998-US6249 30 Mar 1998 PRIORITY INFO: US 1997-829027 31 Mar 1997

DOCUMENT TYPE: Patent LANGUAGE: English

OTHER SOURCE: WPI: 1998-557119 [47]

AB A purified mitochondrial adenylate-kinase

(EC-2.7.4.3) with a given protein sequence is claimed. Also claimed is a nucleic acid encoding the kinase, of given nucleotide sequence, and that hybridizes, under stringent conditions, with the given nucleic acid sequence. The claims also cover a nucleic acid complementary to the given sequence, and a DNA probe that constitutes part of that complementary sequence. Also covered are an expression vector containing the given nucleic acid sequence, a host cell transformed by that vector, and a means of preparing the adenylatekinase by culturing the transformed cell, and recovering the protein. The claims extend to a composition containing the adenylate-kinase, and an antibody, agonist and antagonist of the protein. These are used to treat neurological disorders, cancer and immunological disorders. Also claimed is a means of detecting nucleic acids encoding mitochondrial adenylate-kinase in a sample using the DNA probe, and detecting the hybridization complex. The nucleic acids can also be administered for gene therapy. (63pp)

ANSWER 17 OF 32 HCAPLUS COPYRIGHT 2003 ACS 1998:324881 HCAPLUS ACCESSION NUMBER: 129:39786 DOCUMENT NUMBER: Diabetes-mediating proteins and their therapeutic uses TITLE: Mose, Larsen Peter; Fey, Stephen J.; Nerup, Jorn; INVENTOR(S): Karlsen, Allan E.; Bjerre, Christensen Ulla; Pociot, Flemming; Andersen, Henrik U. Mose Larsen, Peter, Den.; Fey, Stephen J.; Nerup, PATENT ASSIGNEE(S): Jorn; Karlsen, Allan E.; Bjerre Christensen, Ulla; Pociot, Flemming; Andersen, Henrik U. PCT Int. Appl., 145 pp. SOURCE: CODEN: PIXXD2 Patent DOCUMENT TYPE: English LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE \_\_\_\_\_\_ WO 9820124 A2 19980514 WO 1997-IB1627 WO 9820124 A3 19981008 W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG 20010116 JP 1998-513441 19970916 JP 2001500614 Т2 AU 9854070 A1 19980529 AU 1998-54070 19971024 EP 934409 A2 19990811 EP 1997-947839 19971024 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI Т2 JP 1998-520234 19971024 JP 2001503860 20010321 Т2 JP 1998-521182 19971024 JP 2002504806 20020212 KR 2000052802 Α 20000825 KR 1999-703621 19990424 PRIORITY APPLN. INFO.: US 1996-29324P P 19961025 US 1996-30088P P 19961105 US 1996-30186P P 19961105 US 1997-897098 A2 19970718 US 1996-31291P P 19960916 US 1996-29325P P 19961025 WO 1997-IB1114 W 19970916 WO 1997-IB1337 W 19971024 W 19971024 WO 1997-IB1627 Protective and deleterious diabetes-mediating proteins involved in the AB development of diabetes or in the prevention of diabetes development are identified by differential expression during during development

Protective and deleterious diabetes-mediating proteins involved in the development of diabetes or in the prevention of diabetes development are identified by differential expression during during development of diabetes relative to expression in the absence of diabetes development. These proteins are referred to by their position on 10% IEF or NEPHGE 2-dimensional gels. The purified diabetes-mediating proteins are characterized by mol. wt., isoelec. point, and mass spectroscopic characteristics. Galectin-3 (rat and human) and mortalin (mouse and human), two of the identified proteins from pancreatic islets, were also sequenced. Transgenic animals expressing a diabetes-mediating protein, drug screening methods for identifying a test compd. capable of altering the expression of a diabetes-mediating protein, and methods of preventing or ameliorating diabetes by administering a compd. capable of altering the expression of a diabetes-mediating protein are also provided.

L7 ANSWER 18 OF 32 MEDLINE DUPLICATE 5

ACCESSION NUMBER: 1999033072 MEDLINE

DOCUMENT NUMBER: 99033072 PubMed ID: 9813319

TITLE: Identification of a novel adenylate

kinase system in the brain: cloning of

the fourth adenylate kinase.

AUTHOR: Yoneda T; Sato M; Maeda M; Takagi H

CORPORATE SOURCE: First Department of Anatomy, Osaka City University Medical

School, 1-4-3 Asahimachi, Abeno-ku, Osaka-shi, Osaka

545-8585, Japan.

SOURCE: BRAIN RESEARCH. MOLECULAR BRAIN RESEARCH, (1998 Nov 20) 62

(2) 187-95.

Journal code: 8908640. ISSN: 0169-328X.

PUB. COUNTRY: Netherlands

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

OTHER SOURCE: GENBANK-D85036; GENBANK-D87809

ENTRY MONTH: 199902

ENTRY DATE: Entered STN: 19990301

Last Updated on STN: 20000303 Entered Medline: 19990218

AB We identify a novel subtype of adenylate kinase, which is the 4th adenylate kinase (AK4), in the vertebrate.

AK4 mRNA is expressed in the mammalian central nervous system in a region-specific manner from the middle stage of embryogenesis to the adulthood in the rodent. The presence of three isozymes of adenylate kinase (AK1, AK2 and AK3) that maintains the homeostasis of adenine and guanine nucleotide composition has been reported in the vertebrate. Obtained mouse AK4 cDNA is 3667 bp in size. The predicted open reading frame consists of 223 amino acid residues. Rat AK4 cDNA is also obtained, and the predicted open reading frame is the same length as that of the mouse. The predicted rat AK4 molecule shows 97.8% homology with mouse AK4. Rat AK4 protein is distinct from rat AK3,

53.8% homologous with rat AK3, although the adenylate kinase signature and the mitochondrial energy transfer

protein signature are found in both sequences. Interestingly, rat AK4 is 89.2% homologous with the human AK3 over 223 amino acid residues and rat AK3 is 53.7% homologous with the human AK3 indicating that the reported human AK3 actually belongs to the AK4 group (therefore, it should be referred to as human AK4). Although the sequence of AK4 is most similar to that of AK3 among the AK isozymes, its in vivo expression is completely different from AK3; AK4 mRNA is expressed in the pyramidal cells in the hippocampus

(mainly in the subfield CA3), the granular cells in the cerebellum, nasal neuroepithelium and the liver while AK3 mRNA is **expressed** ubiquitously in the body. It is probable that AK4 acts on the specific mechanism of energy metabolism rather than control of the homeostasis of the ADP pool ubiquitously.

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L7 ANSWER 19 OF 32 EMBASE COPYRIGHT 2003 ELSEVIER SCI. B.V.

ACCESSION NUMBER: 97300699 EMBASE

DOCUMENT NUMBER: 1997300699

TITLE: p32 protein, a splicing factor 2-associated protein, is

localized in mitochondrial matrix and is

functionally important in maintaining oxidative

phosphorvlation.

AUTHOR: Muta T.; Kang D.; Kitajima S.; Fujiwara T.; Hamasaki N.

CORPORATE SOURCE: D. Kang, Dept. of Clinical Chem./Lab. Med., Kyushu

University Fac. of Medicine, 3-1-1 Maidashi, Higashi-ku,

Fukuoka 812-82, Japan

Journal of Biological Chemistry, (1997) 272/39 SOURCE:

(24363-24370).

United States

Refs: 44

ISSN: 0021-9258 CODEN: JBCHA3

COUNTRY: DOCUMENT TYPE:

Journal; Article Microbiology

FILE SEGMENT:

004

LANGUAGE: English SUMMARY LANGUAGE: English

Human p32, originally cloned as a splicing factor

2-associated protein, has been reported to interact with a variety of

molecules including human immunodeficiency virus Tat and

complement 1q (C1q). p32 protein is supposed to be in the nucleus and on

the plasma membrane for the association with human

immunodeficiency virus Tat and Clq, respectively. None of the interactions, however, is proven to have a physiological role. To investigate the physiological function of p32, we determined the

intracellular localization of p32. The fractionation of cells, fluorescent immunocytochemistry, and electron microscopic immunostaining show that p32

is exclusively localized in the mitochondrial matrix. We cloned a Saccharomyces cerevisiae homologue of human p32

gene, referred to yeast p30 gene. The yeast p30 protein is also localized

in the mitochondrial matrix. The disruption of the p30 gene

caused the growth retardation of yeast cells in a glycerol medium but not in a glucose medium, i.e. the impairment of the mitochondrial

ATP synthesis. The growth impairment was restored by the introduction of the human p32 cDNA, indicating that p30 is a functional yeast counterpart of human p32. Taken together, both p32 and p30

reside in mitochondrial matrix and play an important role in maintaining mitochondrial oxidative phosphorylation.

ANSWER 20 OF 32 MEDLINE DUPLICATE 6

ACCESSION NUMBER:

1998088919

MEDLINE

DOCUMENT NUMBER:

98088919 PubMed ID: 9428643

TITLE:

Intrinsic nucleoside diphosphate kinase-like activity as a

novel function of 14-3-3 proteins.

AUTHOR:

SOURCE:

Yano M; Mori S; Niwa Y; Inoue M; Kido H

CORPORATE SOURCE:

Division of Enzyme Chemistry, Institute for Enzyme

Research, The University of Tokushima, Japan. FEBS LETTERS, (1997 Dec 15) 419 (2-3) 244-8.

Journal code: 0155157. ISSN: 0014-5793.

PUB. COUNTRY:

Netherlands

DOCUMENT TYPE:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

199801

ENTRY DATE:

Entered STN: 19980206

Last Updated on STN: 19980206 Entered Medline: 19980127

AB 14-3-3 proteins play a role in many cellular functions as molecular chaperone and adapter proteins: they bind to and modulate several proteins involved in cell proliferation and differentiation, and also function ATP-dependently in targeting of precursors to mitochondria. We show here that 14-3-3 purified from a human lymphoblastoma and also its recombinant tau isoform exhibited intrinsic nucleoside diphosphate (NDP) kinase-like activity. 14-3-3 proteins preferentially catalyzed the transfer of the gamma-phosphate group from ATP, dATP or dGTP to all nucleoside diphosphates and this transfer involved acid-labile phosphoenzyme intermediates. They also simultaneously catalyzed the reverse reaction of ATP hydrolysis. These properties of 14-3-3 are similar to those of NDP kinase, but not to those of adenylate kinase.

ANSWER 21 OF 32 SCISEARCH COPYRIGHT 2003 THOMSON ISI

95:187668 SCISEARCH ACCESSION NUMBER:

THE GENUINE ARTICLE: QK489

CONTROL OF CELLULAR RESPIRATION IN-VIVO BY TITLE:

MITOCHONDRIAL OUTER-MEMBRANE AND BY

CREATINE-KINASE - A NEW SPECULATIVE HYPOTHESIS - POSSIBLE

INVOLVEMENT OF MITOCHONDRIAL-CYTOSKELETON

INTERACTIONS

SAKS V A (Reprint); KUZNETSOV A V; KHUCHUA Z A; VASILYEVA AUTHOR:

E V; BELIKOVA J O; KESVATERA T; TIIVEL T

UNIV GRENOBLE 1, PHYSIOL CELLULAIRE CARDIAQUE LAB, BP 53X, CORPORATE SOURCE:

F-38041 GRENOBLE, FRANCE (Reprint); INST CHEM & BIOL PHYS,

BIOENERGET LAB, TALLINN, ESTONIA; CARDIOL RES CTR,

BIOENERGET GRP, MOSCOW 121552, RUSSIA

COUNTRY OF AUTHOR: FRANCE; ESTONIA; RUSSIA

SOURCE:

JOURNAL OF MOLECULAR AND CELLULAR CARDIOLOGY, (JAN 1995)

Vol. 27, No. 1, pp. 625-645.

ISSN: 0022-2828.

DOCUMENT TYPE: General Review; Journal

FILE SEGMENT: LIFE LANGUAGE: ENGLISH REFERENCE COUNT: 119

\*ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS\*

AB The current problems of regulation of myocardial energy metabolism and oxidative phosphorylation in vivo are considered, With this purpose, retarded diffusion of ADP in cardiomyocytes was studied by analysis of elevated apparent K-m for this substrate in regulation of respiration of saponin-skinned cardiac fibers, as compared to isolated mitochondria. Recently published data showing the importance of the outer mitochondrial membrane were compared with new experimental results on the proteolysis of skinned fibers and tissue homogenates. In both cases 10 min incubation and 0.125 mg/ml of trypsin resulted in a decrease of apparent K-m for ADP from 297 +/- 35 and 228 +/-16 to 109 +/- 2 and 36 +/- 16, respectively. Thus, the permeability of the outer mitochondrial membrane for ADP may be controlled by some unknown cytoplasmic protein(s), probably related to the cytoskelton, which are separated from mitochondria during their isolation. The extent of expression of this protein(s) depends on the energy state and type of muscle. Activation of mitochondrial creatine kinase reaction coupled to oxidative phosphorylation overcomes the diffusion difficulties of ADP by amplifying the stimulatory effect of ADP on respiration. It is concluded that both cytoplasmic and mitochondrial creatine kinases, adenylate kinase and cytoplasmic factor controlling outer membrane permeability may

participate in metabolic feedback regulation of respiration in muscle

ANSWER 22 OF 32 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC.

ACCESSION NUMBER: 1995:481319 BIOSIS DOCUMENT NUMBER: PREV199598495619

TITLE: Transfection of a myc gene as a means of generating

infinite life span human fibroblast strains.

McCormick, J. Justin (1); Kohler, Suzanne K.; Maher, AUTHOR(S):

Veronica M.

CORPORATE SOURCE: (1) Carcinogenesis Lab., Fee Hall, Mich. State Univ., East

Lansing, MI 48824-1316 USA

SOURCE: Methods in Cell Science, (1995) Vol. 17, No. 2, pp.

141-148.

ISSN: 1381-5741.

DOCUMENT TYPE: Article LANGUAGE: English

Human fibroblasts in culture have never been found to transform AB spontaneously into immortal cells. In an effort to generate an infinite life span cell strain from foreskin-derived normal diploid fibroblasts, we transfected the cells with a plasmid carrying a v-myc oncogene linked to the neo gene, or with a control vector carrying the neo gene, and selected drug-resistant clones. A clone that expressed the v-myc protein was propagated to the end of its life span, with periodic cryogenic storage of the progeny. The population went into crisis at the same time as cells from the control population and eventually senesced. However, while the cells were senescing, viable-appearing clones were noted. The cells of these clones continued to multiply, very slowly at first but eventually at a faster rate. Analysis showed that these cells have a diploid karyotype that has remained stable throughout more than 200 population doublings since their sibling cells senesced. Molecular analysis showed that the infinite life span cells are, indeed, derived from the cells used for transfection, and that they continue to express the v-myc protein.

ANSWER 23 OF 32 SCISEARCH COPYRIGHT 2003 THOMSON ISI

ACCESSION NUMBER: 94:686537 SCISEARCH

THE GENUINE ARTICLE: PN491

PRIMARY AMINO-ACID-SEQUENCE AND STRUCTURE OF HUMAN TITLE:

PYRUVATE-CARBOXYLASE

WEXLER I D (Reprint); DU Y F; LISGARIS M V; MANDAL S K; AUTHOR:

FREYTAG S O; YANG B S; LIU T C; KWON M; PATEL M S; KERR D

CASE WESTERN RESERVE UNIV, RAINBOW BABIES & CHILDRENS CORPORATE SOURCE:

> HOSP, SCH MED, DEPT BIOCHEM, 2047 ABINGTON RD, CLEVELAND, OH, 44106 (Reprint); CASE WESTERN RESERVE UNIV, UNIV HOSP CLEVELAND, SCH MED, DEPT PEDIAT, CLEVELAND, OH, 44106; HENRY FORD HOSP, MOLEC BIOL RES PROGRAM, DETROIT, MI,

48202

COUNTRY OF AUTHOR:

USA SOURCE:

BIOCHIMICA ET BIOPHYSICA ACTA-MOLECULAR BASIS OF DISEASE,

(21 OCT 1994) Vol. 1227, No. 1-2, pp. 46-52.

ISSN: 0925-4439.

DOCUMENT TYPE:

Article; Journal

FILE SEGMENT:

LIFE

LANGUAGE:

**ENGLISH** 

REFERENCE COUNT:

\*ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS\*

AB Pyruvate carboxylase (PC) (pyruvate:carbon dioxide ligase (ADP-forming), EC 6.4.1.1.), a nuclear-encoded mitochondrial enzyme, catalyzes the conversion of pyruvate to oxaloacetate. We have isolated and characterized cDNAs spanning the entire coding region of human PC. The sequence of human PC has an open reading frame of 3537 nucleotides which encodes for a polypeptide with a length of 1178 amino acids. The identity of the cDNA as PC is confirmed by comparison to PC cDNAs of other species and sequenced peptide fragments of mammalian PC. The M(r) of the full length precursor protein is 129 576 and that of the mature apoprotein is 127 370. RNA blot analysis from a variety of human tissues demonstrates that the highest level of PC mRNA is found in liver corresponding to this tissue's high level of PC activity. Based on homology with other biotin-containing proteins, the ATP, pyruvate, and biotin-binding sites can be identified. One of two patients with documented PC deficiency was found to be missing PC mRNA, further confirming the identity of this cDNA.

ANSWER 24 OF 32 SCISEARCH COPYRIGHT 2003 THOMSON ISI

ACCESSION NUMBER: 92:709675 SCISEARCH

THE GENUINE ARTICLE: KB012

TITLE: VIRAL THYMIDINE KINASES AND THEIR RELATIVES AUTHOR: GENTRY G A (Reprint)

CORPORATE SOURCE: UNIV MISSISSIPPI, MED CTR, DEPT MICROBIOL, JACKSON, MS,

39216 (Reprint)

COUNTRY OF AUTHOR:

PHARMACOLOGY & THERAPEUTICS, (1992) Vol. 54, No. 3, pp. SOURCE:

319-355.

ISSN: 0163-7258.

DOCUMENT TYPE: General Review; Journal

FILE SEGMENT: LIFE LANGUAGE: **ENGLISH** REFERENCE COUNT: 200

\*ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS\*

Thymidine kinases were described for cellular life long before it was AB shown that they could also be encoded by viruses, but the viral thymidine kinase genes were the first to be sequenced. These enzymes have been extraordinarily useful to the researcher, serving first to help label DNA, then to get thymidine analogs incorporated into DNA for therapeutic and other purposes and more recently to move genes from one genome to another. Knowledge of the nucleotide and amino acid sequences of these enzymes has allowed some deductions about their possible three-dimensional structure, as well as the location on the polypeptide of various functions; it has also allowed their classification into two main groups: the herpesviral thymidine/eukaryotic deoxycytidine kinases and the poxviral and cellular thymidine kinases; the relationships of the mitochondrial enzyme are still not clear.

ANSWER 25 OF 32 MEDLINE L7

ACCESSION NUMBER: 90363911 MEDLINE

PubMed ID: 2168054 DOCUMENT NUMBER: 90363911

TITLE: Gene structures of three vertebrate adenylate

kinase isozymes.

AUTHOR: Nakazawa A; Yamada M; Tanaka H; Shahjahan M; Tanabe T

Department of Biochemistry, Yamaguchi University School of CORPORATE SOURCE:

Medicine, Japan.

PROGRESS IN CLINICAL AND BIOLOGICAL RESEARCH, (1990) 344 SOURCE:

495-514.

Journal code: 7605701. ISSN: 0361-7742.

PUB. COUNTRY:

United States

Journal; Article; (JOURNAL ARTICLE) DOCUMENT TYPE:

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199010

ENTRY DATE: Entered STN: 19901109

> Last Updated on STN: 19901109 Entered Medline: 19901003

AB Adenylate kinase is an ubiquitous enzyme which

> contributes to homeostasis of adenine nucleotide composition in the cell. In vertebrates, three isozymes (AK1, AK2, and AK3) are characterized which have distinct distribution in tissues as well as subcellular compartments.

The genetic backgrounds of these adenylate kinase isozymes were analyzed. cDNA clones for AK1 were isolated from

poly(A)+RNA of chicken skeletal muscle. The results of mRNA analysis in

various tissues using the AK1 cDNA indicated that the AK1 gene

expression is regulated both tissue-specifically and

developmentally at the transcriptional level. The AK1 gene was

cloned from chicken and human DNA and characterized.

Both genes were split into seven exons. The intron positions in both genes coincided. cDNA clones for AK2 isolated from bovine liver poly(A)+RNA contained two types. One type (AK2A) encoded the same amino acid sequence as that reported for bovine heart AK2. The other type (AK2B) encoded the same sequence as AK2 except for the COOH terminus. The

mRNA species corresponding to the two cDNA clones were

identified in bovine liver and heart. Both the cDNA sequences were found to direct the active adenylate kinase synthesis in E. coli. The AK2 gene was cloned and characterized. It consisted of seven exons and six introns. From genomic structure analysis, the two cDNA species were shown to be derived from a single gene by the alternative splicing mechanism. Three types of cDNA clones for AK3 were isolated from bovine liver poly(A)+RNA, which contained the common AK3-coding region and different 3' portions. No NH2-terminal presequence of mitochondrial targeting was identified in AK3 from the sequencing and expression analyses of the cDNA. Upon expression of the cDNA sequence in E. coli, AK3 protein was recovered in the periplasmic space of the bacteria, indicating that AK3 without presequence was exported through the inner bacterial membrane as it is imported through the mitochondrial membranes. Internal targeting signals may be responsible for the translocation process. AK3 gene was cloned and partially characterized. It is split into at least five exons. The comparisons of amino acid sequences and genomic structure of three isozymes revealed that a segment corresponding to either exon 5 of the AK2 gene or a part of exon 3 of the AK3 gene is missing in the AK1 gene. Phylogenetic analysis suggested that AK1, a shorter molecule, would have been separated from a longer molecule very early in evolution of adenylate kinase. (ABSTRACT TRUNCATED AT 400 WORDS)

L7 ANSWER 26 OF 32 MEDLINE

ACCESSION NUMBER: 90037053 MEDLINE

DOCUMENT NUMBER: 90037053 PubMed ID: 2478555

TITLE: Cloning and characterization of cDNA for

mitochondrial GTP: AMP phosphotransferase of bovine

liver.

AUTHOR: Yamada M; Shahjahan M; Tanabe T; Kishi F; Nakazawa A

CORPORATE SOURCE: Department of Biochemistry, Yamaguchi University School of

Medicine, Japan.

SOURCE: JOURNAL OF BIOLOGICAL CHEMISTRY, (1989 Nov 15) 264 (32)

19192-9.

Journal code: 2985121R. ISSN: 0021-9258.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals OTHER SOURCE: GENBANK-M25757

ENTRY MONTH: 198912

ENTRY DATE: Entered STN: 19900328

Last Updated on STN: 19960129 Entered Medline: 19891215

AΒ Three different types of cDNA clones for mitochondrial GTP:AMP phosphotransferase (AK3) were isolated from a cDNA library of bovine liver poly(A) + RNA. Nucleotide sequencing revealed that each of these clones consisted of a common 5'-untranslated region, a common AK3-coding sequence and a 3'-untranslated region with different sizes. By Northern blot analysis, three species of AK3 mRNA apparently corresponding to the isolated cDNA clones were detected, which would be a result of varying terminations and polyadenylations of the primary transcript. From comparison of the size of the product synthesized in vitro from the message directed by the isolated cDNA with that of the purified AK3 protein, AK3 appeared to have no cleavable NH2-terminal sequence as found in other mitochondrial proteins. The AK3 cDNA was expressed in Escherichia coli, which resulted in complementation of an adenylate kinase mutation of The AK3 product was exported to the periplasmic space through the bacterial inner membrane. The possible involvement of the NH2-terminal sequence of the protein in targeting to the

## mitochondrial matrix was discussed.

ANSWER 27 OF 32 EMBASE COPYRIGHT 2003 ELSEVIER SCI. B.V.DUPLICATE 7

ACCESSION NUMBER: 83053333 EMBASE

DOCUMENT NUMBER: 1983053333

Adenosine triphosphate-adenosine-5'-monophosphate TITLE:

phosphotransferase from normal human liver

mitochondria. Isolation, chemical properties, and

immunochemical comparison with Duchenne dystrophic serum

aberrant adenylate kinase.

Hamada M.; Sumida M.; Okuda H.; et al. AUTHOR:

CORPORATE SOURCE: Dep. Hyg., Ehime Univ. Sch. Med., Shigenobu cho, Onsen gun,

Ehime 791-02, Japan

SOURCE: Journal of Biological Chemistry, (1982) 257/21

> (13120-13128). CODEN: JBCHA3 United States

COUNTRY: DOCUMENT TYPE: Journal

FILE SEGMENT: 029 Clinical Biochemistry

> 800 Neurology and Neurosurgery

LANGUAGE: English

AB

Adenylate kinase has been purified approximately 1360-fold to a final specific activity of 280 .mu.mol of ATP formed min-1xmg-1 of protein at 30.degree.C from normal human liver

mitochondria. The purity of the final preparation was evaluated by studies with polyacrylamide gel electrophoresis and sodium dodecyl sulfate-polyacrylamide gel electrophoresis and by sedimentation studies. The purified enzyme catalyzes transphosphorylation reactions between adenosine triphosphate (ATP) and adenosine monophosphate (AMP). ATP and adenosine-5'-thiophosphate, ATP and adenosine monophosphate-3'pyrophosphate, adenosine-s'-(3-thio)triphosphate and AMP. The nearly constant ratios of these activities throughout the purification scheme suggest that all are catalyzed by the same enzyme. The purified enzyme has a molecular weight of 25,200 by sedimentation equilibrium with the use of a partial specific volume of 0.73 mlxg-1 calculated from amino acid analysis. This purified enzyme was also found to be a single polypeptide with a molecular weight of 26,500 by sodium dodecyl sulfate-polyacrylamide gel electrophoresis. From amino acid analysis, a calculated minimum molecular weight of 26,349 was obtained. Initial velocity studies revealed a narrow specificity for adenine nucleotides. The Kd' values for MgATP2and MgATP2-.gamma.S1 were 0.12 and 0.57 .mu.M with Vmax.forward values of 1.04 (.+-.0.04) x103 and 7.02x102 .mu.mol x min-1 x mg-1, respectively. For the monophosphate acceptor, Kd' values of 0.56 and 186 .mu.M were measured for 5'-AMP2- and AMP2-.alpha.S, respectively. The Kd' for MgADP1- and ADP3- were 0.53 and 0.17 .mu.M with a Vmax.reverse of  $6.40(.+-.0.03)\times102$ .mu.molxmin-lxmg-1 of protein. The steady state kinetics, at pH 7.4, 30.degree.C, and essentially fixed .DELTA./2 of 0.16-0.18, of this enzyme seem to be adequately expressed by a random quasi-equilibrium type of mechanism with a rate-limiting step largely at the interconversion of the ternary complexes, as shown in rabbit muscle, calf muscle, and calf liver adenylate kinase. It would appear that normal

human liver mitochondrial adenylate

kinase largely favors the forward reaction (ADP formation). A specific anti-liver enzyme antibody obtained from rabbit serum inhibited the purified liver mitochondrial enzyme activity, but not the purified human muscle enzyme, nor the aberrant adenylate kinase from Duchenne dystrophic serum.

ANSWER 28 OF 32 MEDLINE **DUPLICATE 8** 

ACCESSION NUMBER: 82003493 MEDLINE

DOCUMENT NUMBER: 82003493 PubMed ID: 6944169

TITLE: Characterization of the Philadelphia chromosome by gene mapping.

Geurts van Kessel A H; ten Brinke H; Boere W A; den Boer W AUTHOR:

C; de Groot P G; Hagemeijer A; Meera Khan P; Pearson P L

CYTOGENETICS AND CELL GENETICS, (1981) 30 (2) 83-91. SOURCE:

Journal code: 0367735. ISSN: 0301-0171.

PUB. COUNTRY: Switzerland

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 198111

Entered STN: 19900316 ENTRY DATE:

Last Updated on STN: 19900316 Entered Medline: 19811122

AB Chinese hamster X human and mouse X human somatic cell hybrid lines were obtained using circulating leucocytes from six chronic myeloid leukemia patients. All six patients carried the Ph1 translocation, t(9q+;22q-), characteristic of chronic myeloid leukemia, in their dividing immature granulocytes. Analysis of independent hybrid clones yielded the following results: 1. The chromosome 9 markers, soluble aconitase and adenylate kinase-1, segregated with the 9q+ derivative. The latter marker has previously been localized to 9q34. 2. The chromosome 22 markers, mitochondrial aconitase, N-acetyl-alpha-D-galactosaminidase, and arylsulfatase-A, also segregated with the 9q+ derivative. Mitochondrial aconitase has recently been assigned to 22q11 leads to 22q13. No evidence was obtained either for reciprocity of the translocation or for variations in breakpoints in different patients. The results reported in this paper provisionally assign the gene for mitochondrial aconitase to a region distal to the breakpoint in 22q11.

ANSWER 29 OF 32 LIFESCI COPYRIGHT 2003 CSA L7

81:24127 LIFESCI ACCESSION NUMBER:

TITLE: Characterization of the Philadelphia Chromosome by Gene

Van Kessel, A.H.M.G.; Ten Brinke, H.; Boere, W.A.M.; Den AUTHOR:

Boer, W.C.; De Groot, P.G.; Hagemeijer, A.; Meera Khan, P.;

Pearson, P.L.

CORPORATE SOURCE: Dept. Cell Biol. Genet., Erasmus Univ., P.O. Box 1738, 3000

DR Rotterdam, Netherland

CYTOGENET. CELL GENET., (1981) vol. 30, no. 2, pp. 83-91. SOURCE:

DOCUMENT TYPE: Journal

FILE SEGMENT:

LANGUAGE: English SUMMARY LANGUAGE: English

Chinese hamster x human and mouse x human somatic cell hybrid lines were obtained using circulating leucocytes from six chronic myeloid leukemia patients. All six patients carried the Ph super(1) translocation, t(9q+;22q-), characteristic of chronic myeloid leukemia, in their dividing immature granulocytes. Analysis of independent hybrid clones yielded the following results: 1. The chromosome 9 markers, soluble aconitase and adenylate kinase-1, segregated with the 9q+ derivative. The latter marker has previously been localized to 9q34. 2. The chromosome 22 markers, mitochondrial aconitase, N-acetyl- alpha -D-galactosaminidase, and arylsulfatase-A, also segregated with the 9q+ derivative. Mitochondrial aconitase has recently been assigned to 22q11 arrow right 22q13. No evidence was obtained either for reciprocity of the translocation or for variations in breakpoints in different patients.

L7 ANSWER 30 OF 32 MEDLINE

MEDLINE ACCESSION NUMBER: 79194246

PubMed ID: 36399 DOCUMENT NUMBER: 79194246

TITLE: Cytosolic phosphorylation potential.

AUTHOR: Veech R L; Lawson J W; Cornell N W; Krebs H A

SOURCE: JOURNAL OF BIOLOGICAL CHEMISTRY, (1979 Jul 25) 254 (14)

6538-47.

Journal code: 2985121R. ISSN: 0021-9258.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 197909

ENTRY DATE: Entered STN: 19900315

Last Updated on STN: 19990129 Entered Medline: 19790901

The tissue contents of the reactants of the myokinase (EC 2.7.4.3) and the AB combined glyceraldehyde-3-phophate dehydrogenase (EC 1.1.1.29)-3phosphoglycerate kinase (EC 2.7.2.3) reactions were measured in rapidly inactivated samples of human blood and rat brain, muscle, and liver. The tissue contents of the reactants of the creatine kinase (EC 2.7.3.2) reaction were measured in rat brain and muscle. In vitro the value of the expression: KG+G = [sigma3PG] . [sigmaATP] . [sigmalactate] KLDH = [sigmaHAP]/22] . [sigmaADP][sigmaPi] . [sigmaRUVATE] (1) was found to be  $0.725 \times 10(7) \text{ M-1}$  at I = 0.25, T = 38 degrees C, and free [Mq2+] = 0.15 mM and the value measured in vivo in red cell was 0.699  $\times$  10(7) M-1. The value of the expression KMYK = ([sigma ATP] [sigma AMP]/[ADP2]) measured under the above conditions and at pH 7.2 was found to be 0.744 while the value found in red cell was 0.784 +/- 0.037. These reactions, therefore, appear to be in a state of near-equilibrium in the red cell and the measured tissue contents of ATP and ADP, which are common reactants in both reactions, approximate closely the activity of these reactants in vivo. In brain and muscle, the value of KG + G/KLDH calculated from the measured tissue contents of the reactants was a factor of 20 or more lower than that expected at equilibrium as was the measured value of the expression: KCK = [sigma ATP] [sigma creatine] divided by [sigma ADP] [sigma creatine-P] [H+] (2) Substitution of calculated free [sigma ADP] values in the expression of KG + G/KLDH gave values of  $0.83 +/- 0.19 \times 10(7)$  M-1 for brain and muscle, respectively, which agreed well with the value of 1.65 x 10(7) M-1 measured in vitro at I = 0.25, free [Mq2+] = 1 mM, T = 38 degrees C. agreement between two highly active enzyme systems in the same compartment is taken as evidence of the existence of near-equilibrium in both these systems and suggests that free cytosolic [sigma ADP] is probably 20-fold lower than measured cell ADP content in mitochondrial-containing tissues.

L7 ANSWER 31 OF 32 MEDLINE DUPLICATE 9

ACCESSION NUMBER: 77221531 MEDLINE

DOCUMENT NUMBER: 77221531 PubMed ID: 195572

TITLE: Adenylate kinase 2, a mitochondrial enzyme.

AUTHOR: Bruns G A; Regina V M

SOURCE: BIOCHEMICAL GENETICS, (1977 Jun) 15 (5-6) 477-86.

Journal code: 0126611. ISSN: 0006-2928.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 197709

ENTRY DATE: Entered STN: 19900314

Last Updated on STN: 19970203 Entered Medline: 19770902

AB The subcellular compartmentalization of the isoenzymes of ATP:AMP phosphotransferase (adenylate kinase) was analyzed in

HeLa cells, RAG cells, and RAG-human hybrids that express human AK-2. In HeLa cells and in the hybrids, human AK-2 was present in a mitochemical fraction prepared from cell extracts and in mitochondria purified by density gradient centrifugation. Human AK-1 was, as expected, distributed in the soluble cytoplasmic fraction of the cells. The rodent isozymes which are homologous to human AK-1 and AK-2 have been determined.

L7 ANSWER 32 OF 32 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC.

ACCESSION NUMBER: 1977:15881 BIOSIS

DOCUMENT NUMBER: BR13:15881

TITLE: ASSIGNMENT OF HUMAN GENES BETA GLUCURONIDASE TO

CHROMOSOME 7 ADENYLATE KINASE 1 TO 9 A 2ND ENZYME WITH ENOLASE ACTIVITY TO 12 AND MITOCHONDRIAL ISO CITRATE DEHYDROGENASE TO 15.

AUTHOR(S): GRZESCHIK K-H

SOURCE: Cytogenet. Cell Genet., (1976) 16 (1-5), 142-148.

CODEN: CGCGBR. ISSN: 0301-0171.

FILE SEGMENT: BR; OLD LANGUAGE: Unavailable

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L8 8 "HMAK"

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PROCESSING COMPLETED FOR L8

L9 3 DUP REM L8 (5 DUPLICATES REMOVED)

=> d 1-3 ibib ab

L9 ANSWER 1 OF 3 MEDLINE DUPLICATE 1

ACCESSION NUMBER: 2002485451 MEDLINE

DOCUMENT NUMBER: 22217966 PubMed ID: 12084720

TITLE: Identification of human male germ cell-associated kinase, a

kinase transcriptionally activated by androgen in prostate

cancer cells.

AUTHOR: Xia Liang; Robinson Dan; Ma Ai-Hong; Chen Hua-Chien; Wu

Frederick; Qiu Yun; Kung Hsing-Jien

CORPORATE SOURCE: Department of Biological Chemistry, School of Medicine,

University of California, Davis, California 95616, USA.

CONTRACT NUMBER: CA39207 (NCI)

CA57179 (NCI) CA82073 (NCI) DK52659 (NIDDK)

SOURCE: JOURNAL OF BIOLOGICAL CHEMISTRY, (2002 Sep 20) 277 (38)

35422-33.

Journal code: 2985121R. ISSN: 0021-9258.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals OTHER SOURCE: GENBANK-AF505623

ENTRY MONTH: 200210

ENTRY DATE: Entered STN: 20020926

Last Updated on STN: 20030105 Entered Medline: 20021024

AB Androgen is involved in both normal development and malignant transformation of prostate cells. The signal transduction pathways associated with these processes are not well understood. Using a novel kinase display approach, we have identified a protein kinase, human male germ cell-associated kinase (hmak), which is transcriptionally

induced by the androgenic hormone 5alpha-dihydrotestosterone (DHT). kinetics of induction is rapid and dose-dependent, and the induction is not blocked by cycloheximide treatment. Real time reverse transcription-PCR studies demonstrated a 9-fold induction of hMAK by 10 nm DHT at 24 h post-stimulation. The expression levels of hMAK in prostate cancer cell lines are in general higher than those of normal prostate epithelial cells. A reverse transcription-PCR product encompassing the entire hMAK open reading frame was isolated. The results from sequencing analysis showed that the hMAK protein is 623 amino acids in length and contains a kinase catalytic domain at its N terminus, followed by a proline/glutamine-rich domain. The catalytic domain of this kinase contains sequence motifs related to both the cyclin-dependent kinase and the mitogen-activated protein kinase families. When expressed in COS1 cells, hMAK is kinase-active as demonstrated by autophosphorylation and phosphorylation of exogenous substrate and is localized in the nucleus. A 3.7-kilobase pair promoter of the hMAK locus was isolated from a human genomic DNA bacterial artificial chromosome clone and was shown to be activated by DHT. This activation can be blocked by an anti-androgen drug bicalutamide (Casodex), implicating the involvement of androgen receptor in this process. Taken together, these data suggest that hMAK is a protein kinase targeted by androgen that may participate in androgen-mediated signaling in prostate cancer cells.

L9 ANSWER 2 OF 3 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC.

ACCESSION NUMBER: 2000:290852 BIOSIS DOCUMENT NUMBER: PREV200000290852

TITLE: Mitochondrial adenylate kinase.

AUTHOR(S): Hillman, Jennifer L.; Shah, Purvi

ASSIGNEE: Incyte Pharmaceuticals, Inc.

PATENT INFORMATION: US 6001624 December 14, 1999

TATENT INFORMATION. 05 0001024 December 14, 1999

SOURCE: Official Gazette of the United States Patent and Trademark Office Patents, (Dec. 14, 1999) Vol. 1229, No. 2, pp. No

pagination. e-file. ISSN: 0098-1133.

DOCUMENT TYPE: Patent LANGUAGE: English

AB The present invention provides a human mitochondrial adenylate kinase ( HMAK) and polynucleotides which encode HMAK. The invention also provides expression vectors, host cells, agonists, antisense molecules, antibodies, or antagonists. The invention also provides methods for treating disorders associated with expression of HMAK.

L9 ANSWER 3 OF 3 BIOTECHDS COPYRIGHT 2003 THOMSON DERWENT AND ISI

ACCESSION NUMBER: 1999-00127 BIOTECHDS

TITLE: Human mitochondrial adenylate-kinase, HMAK;

sense, antisense sequence, antibody, agonist and

antagonist used for cancer, neurological and immunological

disorder diagnosis and therapy

AUTHOR: Hillman J L; Shah P

PATENT ASSIGNEE: Incyte-Pharm.

LOCATION: Palo Alto, CA, USA.
PATENT INFO: WO 9844124 8 Oct 1998

APPLICATION INFO: WO 1998-US6249 30 Mar 1998 PRIORITY INFO: US 1997-829027 31 Mar 1997

DOCUMENT TYPE: Patent LANGUAGE: English

OTHER SOURCE: WPI: 1998-557119 [47]

AB A purified mitochondrial adenylate-kinase (EC-2.7.4.3) with a given protein sequence is claimed. Also claimed is a nucleic acid encoding the kinase, of given nucleotide sequence, and that hybridizes, under

stringent conditions, with the given nucleic acid sequence. The claims also cover a nucleic acid complementary to the given sequence, and a DNA probe that constitutes part of that complementary sequence. Also covered are an expression vector containing the given nucleic acid sequence, a host cell transformed by that vector, and a means of preparing the adenylate-kinase by culturing the transformed cell, and recovering the protein. The claims extend to a composition containing the adenylate-kinase, and an antibody, agonist and antagonist of the protein. These are used to treat neurological disorders, cancer and immunological disorders. Also claimed is a means of detecting nucleic acids encoding mitochondrial adenylate-kinase in a sample using the DNA probe, and detecting the hybridization complex. The nucleic acids can also be administered for gene therapy. (63pp)

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E1
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E2
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E3
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E4
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E7
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     FILE 'MEDLINE, EMBASE, BIOSIS, BIOTECHDS, SCISEARCH, HCAPLUS, NTIS,
     LIFESCI' ENTERED AT 12:28:34 ON 06 MAY 2003
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           2615 S HUMAN AND L1
L3
         613747 S MITOCHONDR?
            255 S L2 AND L3
L4
L5
        5929362 S CLON? OR EXPRESS? OR RECOMBINANT
             59 S L4 AND L5
L6
             32 DUP REM L6 (27 DUPLICATES REMOVED)
L7
L8
              8 S "HMAK"
L9
              3 DUP REM L8 (5 DUPLICATES REMOVED)
                E HILLMAN J. L/AU
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     ANSWER 1 OF 1 BIOTECHDS COPYRIGHT 2003 THOMSON DERWENT AND ISI
L11
AN
      1999-00127 BIOTECHDS
ΤI
      Human mitochondrial adenylate-
     kinase, HMAK;
         sense, antisense sequence, antibody, agonist and antagonist used for
         cancer, neurological and immunological disorder diagnosis and therapy
·AU
      Hillman J L; Shah P
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Incyte-Pharm.
PA
LO
      Palo Alto, CA, USA.
      WO 9844124 8 Oct 1998
PΙ
      WO 1998-US6249 30 Mar 1998
ΑI
     US 1997-829027 31 Mar 1997
PRAI
DT
      Patent
LA
      English
      WPI: 1998-557119 [47]
os
AB
      A purified mitochondrial adenylate-kinase
      (EC-2.7.4.3) with a given protein sequence is claimed. Also claimed is a
      nucleic acid encoding the kinase, of given nucleotide sequence, and that
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      sequence. The claims also cover a nucleic acid complementary to the
      given sequence, and a DNA probe that constitutes part of that
      complementary sequence. Also covered are an expression vector containing
      the given nucleic acid sequence, a host cell transformed by that vector,
      and a means of preparing the adenylate-kinase by
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      using the DNA probe, and detecting the hybridization complex.
      nucleic acids can also be administered for gene therapy. (63pp)
CC
      D PHARMACEUTICALS; D3 Peptides and Proteins; D PHARMACEUTICALS; D7
      Clinical Genetic Techniques; D PHARMACEUTICALS; D6 Antibodies; A GENETIC
      ENGINEERING AND FERMENTATION; Al Nucleic Acid Technology
     HUMAN MITOCHONDRIA RECOMBINANT ADENYLATE-
CT
     KINASE PREP., VECTOR EXPRESSION IN HOST CELL, DNA PROBE,
     ANTIBODY, AGONIST, ANTAGONIST, SENSE, ANTISENSE SEQUENCE, APPL. CANCER,
     NEUROLOGICAL DISORDER, IMMUNOLOGICAL DISORDER THERAPY, GENE THERAPY,
      DIAGNOSIS ANIMAL MAMMAL ENZYME EC-2.7.4.3 DNA SEQUENCE PROTEIN SEQUENCE
      GENE TRANSFER HYBRIDIZATION CLONING TUMOR (VOL.18, NO.1)
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E11
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L12
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=> df his

DF IS NOT A RECOGNIZED COMMAND

The previous command name entered was not recognized by the system. For a list of commands available to you in the current file, enter "HELP COMMANDS" at an arrow prompt (=>).

=> d his

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FILE 'MEDLINE, EMBASE, BIOSIS, BIOTECHDS, SCISEARCH, HCAPLUS, NTIS,
     LIFESCI' ENTERED AT 12:28:34 ON 06 MAY 2003
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.L2
           2615 S HUMAN AND L1
L3
         613747 S MITOCHONDR?
L4
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L5
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             59 S L4 AND L5
L6
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             32 DUP REM L6 (27 DUPLICATES REMOVED)
              8 S "HMAK"
L8
              3 DUP REM L8 (5 DUPLICATES REMOVED)
L9
                E HILLMAN J L/AU
            454 S E3
L10
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                E SHAH P/AU
T.12
           1409 S E3
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      ANSWER 1 OF 1 BIOTECHDS COPYRIGHT 2003 THOMSON DERWENT AND ISI
ACCESSION NUMBER: 1999-00127 BIOTECHDS
                  Human mitochondrial adenylate-
                  kinase, HMAK;
                     sense, antisense sequence, antibody, agonist and
                     antagonist used for cancer, neurological and immunological
                     disorder diagnosis and therapy
AUTHOR:
                  Hillman J L; Shah P
                  Incyte-Pharm.
PATENT ASSIGNEE:
                  Palo Alto, CA, USA.
LOCATION:
PATENT INFO:
                  WO 9844124 8 Oct 1998
APPLICATION INFO: WO 1998-US6249 30 Mar 1998
                  US 1997-829027 31 Mar 1997
PRIORITY INFO:
DOCUMENT TYPE:
                  Patent
                  English
LANGUAGE:
OTHER SOURCE:
                  WPI: 1998-557119 [47]
AB
      A purified mitochondrial adenylate-kinase
      (EC-2.7.4.3) with a given protein sequence is claimed. Also claimed is a
      nucleic acid encoding the kinase, of given nucleotide sequence, and that
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                                                                   The claims
      extend to a composition containing the adenylate-kinase
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      nucleic acids can also be administered for gene therapy. (63pp)
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L2	2615	S HUMAN AND L1
L3	613747	S MITOCHONDR?
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L5	5929362	S CLON? OR EXPRESS? OR RECOMBINANT
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L7	32	DUP REM L6 (27 DUPLICATES REMOVED)
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